PATENT SPECIFICATION

NO DRAWINGS



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COMPLETE SPECIFICATION

2-Amino-1-(3,4-Methylenedioxyphenyl)-Propane Isomers and an Ataractic preparation containing 2-Amino-1-(3,4-Methylenedioxyphenyl)-Propane

We, SMITH KLINE & FRENCH LABORAtories, a Corporation organized under the Laws of the State of Delaware, one of the United States of America, of 1530, Spring 5 Garden Street, City of Philadelphia, Pennsylvania, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to novel isomers of 2 - amino - 1 - (3,4 - methylenedioxyphenol) propane, and to a medicinal preparation hav-ing ataractic activity.

Prior to the present invention the important advances in the treatment of mentally deranged have largely been in the excited group of patients through the use of central nervous system depressant compounds commonly referred to as tranquilizers. A large proportion of the population of mental hospitals, however, consists of depressed patients whose conditions generally are either not responsive to tranquilizers or aggravated by the use of these drugs. The need of a safe, effective composition for use in this area has been great.

The preparation in accordance with this 30 invention contains 2-amino-1-(3,4-methylenedioxyphenyl)-propane and is very useful in treating various depressive states of psychotic patients due to having an unusual differ-ential in its acitivity. It, surprisingly for 35 a central nervous stimulant, provides a strong conditioned response block in animals. In the treatment of severely depressed psycho-tics, it induces ataraxia without any substantial amount of the sympathomimetic action 40 found in closely related compounds such as amphetamine. This preparation has a low incidence of side effects in a dosage range where preparations containing closely related compounds such as 2-amino-1-phenylpro-panes produce severe side effects such as jitteriness, excessive stimulation or increased tension.

More specifically, the preparation of this invention is in a dosage unit form and comprises from about 15 mg. to about 150 mg., and preferably from about 25 mg, to about 100 mg., of 2-amino-1-(3,4-methylenedioxyphenyl)-propane or a non-toxic acid solution salt thereof and a pharmaceutical carrier.

The d- or 1-isomer of 2-amino-1-(3,4methylenedioxyphenyl)-propane or a nontoxic salt thereof can be substituted advantageously for the racemic mixture. Where the term 2-amino-1-(3,4-methylenedioxyphenyl)-propane is employed without any indication as to the d-, L or racemic form, it is intended herein and in the claims to cover the individual d- and l-isomers as well as mixtures thereof.

The l-isomer is advantageous since it also is an effective anorexic agent and, hence, its employment is advantageous where it is desired to curb the appetite.

The active d-isomer is prepared by dissolving the racemic hydrochloride salt in water, neutralizing with an inorganic base, for example, sodium hydroxide, and extracting into an organic solvent such as ether or benzene. d-Tartaric acid is added to separate the d-tartrate sait. Recrystallization from alcohol such as isopropanol or aqueous isopropanol gives the pure d-isomer as the d-tartrate with an optical rotation of 29.4° (2% in water). The d-base in hexane has a rotation of 24.6° (1%). If desired, the hydrochloride salt may be regenerated from the active base by treating an ether or hexane solution with anhydrous hydrogen chloride gas. The 1-base is similarly prepared.

Preferably the hydrochloric salt of the 2 - amino - 1 - (3.4 - methylenedioxyphenyl) -

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	propane is used, however, either the base itself or a non-toxic pharmaceutically acceptable acid addition salt of the base may be used, such as the salt derived from sulfuric,	ture after seeding. A thick precipitate separates. After filtration, the solid tartrate is recrystallized several times from isopropanol to white crystals of d-2-amino-1-(3,4-	65
5	nitric, phosphoric, citric, acetic, lactic, sali- cylic, tartaric, ethanedisulfonic, sulfamic, acetylsalicylic, succinic, fumaric, maleic, hyd- robromic, or benzoic acid. The salts are conveniently prepared by reacting the free	methylenedioxyphenyl)-propane d-tartrate, m.p. 145—146° C., [a] ²⁵ and 29.44° (1% H ₂ O). The free d-base is regenerated and taken into hexane, [a] ²⁵ +24.6°. The free d-base is reconverted to the hydrochloride	70
10	base with either a stoichiometric amount or an excess of the desired acid in a suitable sol-	salt with gaseous hydrogen chloride, m.p. 185 —187° C.	75
	vent such as ethanol, ether, ethyl acetate, acetone, water or various combinations of solvents.	The mother filtrate is evaporated to give 22 g. of the 1-2-amino-1-(3,4-methylenedioxyphenyl)-propane d-tartrate, m.p. 125—130° C. After converting a portion to the base	
15	The lower part of the dasage range of the 2 - amino - 1 - (3,4 - methylenedioxyphenyl) - propane of from about 15 mg. to about 25 mg. is aimed at child medication and at	in hexane, the specific rotation of this sample is -11.5° C. The remainder of the tartrate is recrystallized from aqueous ethanol to pure	80
20	parenteral preparations. For oral use with a solid carrier the preparation for adults would advantageously contain from about 25 mg.	white crystals of <i>l</i> -base <i>d</i> -tartrate, m.p. 129—137.° C., $[z]^{25}$ —28.5° (1% H_2O).	
	to about 75 mg. of the active propane com- pound. It a sustained release (i.e. having a release over a period of about 12 hours) is	EXAMPLE 2 dl - 2 - Amino - 1 - (3,4 - methylene - dioxyphenyl)-propane Hadrophorida 25 mg	85
25	used, the above dosage ranges can be tripled. The pharmaceutical carrier may be, for example, either a solid or a liquid. Exemplary of solid carriers are tale, corn starch,	Hydrochloride - 25 mg. Lactose 230 mg. Starch 45 mg. The above ingredients were thoroughly	90
30	lactose, ethylcellulose, magnesium stearate, agar, pectin, stearic acid, gelatin and acacia. Exemplany of liquid carriers are water, pea-	mixed, granulated using a 10% gelatin solu- tion and compressed into tablets using an admixture of talc-stearic acid as a lubricant.	
	nut oil, olive oil and sesame oil. Solid carriers are preferred. A wide variety of pharmaceutical forms	EXAMPLE 3 dl - 2 - Amino - 1 - (3,4 - methylene -	95
35	can be employed. Thus, if a solid carrier is used, the preparation can be tabletted or placed in a hard gelatin capsule. If a liquid	dioxyphenyl)-propane Maleate 75 mg. Lactose 225 mg.	
40	carrier is used, the preparation may be in the form of a soft gelatin capsule or placed in an ampule. The amount of carrier will vary widely but preferably will be from about 25	The above ingredients were thoroughly mixed, granulated using a 50% sucrose solution and compressed into tablets using an admixture of 7% starch and 1% magnesium stearate based on tablet weight.	100
45	mg. to about 1 gm. The preparation of this invention may be administered internally in an amount to produce the comparison of the property of the product o	EXAMPLE 4 d - 2 - Amino - 1 - (3,4 - methylene -	105
45	duce ataraxia in depressed psychotic patients. The administration may be orally or parenterally preferably employing the above described preparation. In this method it is preferred	dioxyphenyl)-propane Hydrochloride - 50 mg. Lactose 150 mg.	•••
50	to administer from about 60 mg. to about 350 mg. and advantageously about 75 mg. to about 320 mg. of 2-amino-1-(3,4-methylene-dioxyphenyl)-propane or a salt thereof daily.	Starch 50 mg. The above ingredients were thoroughly mixed, granulated using a 10% gelatin solution and compressed into scored tablets.	110
55	preferably administering equal doses three or four times daily. In the treatment of children somewhat lower dosages are used	Example 5	
	depending largely on the age and weight of the child. Such doses may be individually determined by the physician but will ordin- arily be about half the adult dosage.	dl - 2 - Amino - 1 - (3,4 - methylene - dioxyphenyl)-propane Hydrochloride - 300.00 gm. Lactose	115
60	Example 1	(200 mesh) - 2820.00 gm. Magnesium stearate 60.00 gm.	120
	A solution of 35.8 g. (0.2 mole) of 2-amino- 1-(3.4-methylenedioxyphenyl)-propane and 30 g. of d-tartaric acid in 600 ml. of 75% iso- propanol is allowed to stand at room tempera-	The powders are mixed, screened and filled into No. 2 hard gelatin capsules (12,000 capsules at 25 mg).	

	EXAMPLE 6 1 - 2 - Amino 1 - (3,4 - methylene dioxyphenyl)-propane Sulfate 75 mg	dioxyphenyl)-propane	· 50
5	Sulfate 75 mg. Peanut oil - 225 mg. The ingredients are mixed to a thick slurry and filled into a soft gelatin capsule.	Hydrochloride - 2.0 w/v Sodium chloride - 0.375 w/v Water for injection, U.S.P., q.s. ad 100 %	55
10		The solid ingredients are dissolved in part	
	Hydrochloride - 100 mg. Hydrogenated castor oil - 100 mg.	WHAT WE CLAIM IS:—	60
15	The chemical is imbedded in the hydro- genated castor oil by melting the latter, mix- ing in the chemical and solidifying. After-	prising a pharmaceutical carrier and a 2-	
20	comminuting and screening through a Num- ber 10 screen, the powder is granulated with a small amount of starch to produce sustained release granules.	propane or its non-toxic acid addition salts. 2. The preparation claimed in Claim 1 in which the dosage unit form is a capsule	65
	dl - 2 - Amino - 1 - (3,4 - methylene - dioxyphenyl)-propane Hydrochloride - 50 mg	which the dosage unit form is a tablet. 4. The preparation claimed in any of Claims.	70
25	Talc 15 mg. The above ingredients are mixed and	1 to 3 in which the 2-amino-1-(3,4-methylene-dioxyphenyl)-propane is in the racemic form. 5. The preparation claimed in any of Claims 1 to 3 in which the 2-amino-1-(3,4-methylene-dioxynday).	75
30	granulated with a gelatin solution, dried, screened and compressed into cylindrical, flat faced tablets. The sustained release granules are added to the die and compressed onto the previously formed tablets.	dioxyphenyl)-propane is in the dextro isomer. 6. The preparation claimed in any of Claims 1 to 3 in which the 2-amino-1-(3,4-methylenedioxyphenyl)-propane is the levo isomer.	75
	EXAMPLE 8 d - 2 - Amino - 1 - (3,4 - methylene - dioxyphenyl)-propane	7. The preparation claimed in any of the preceding claims in which the 2-amino-1-(3,4-methylenedioxyphenyl)-propage of its per-	80
35	Lactose 245 mg. Magnesium stearate 5 mg.	toxic acid addition salts are present in an amount of from about 15 mg to about 150 mg. 8. The preparation claimed in any of Claims 1 to 6 in which the 2-amino-1-(3,4-methyleradia-methyl	85
	The powders are mixed, screened and filled into a Number 2 hard gelatin capsule.	methylenedioxyphenyl)-propane or its non- toxic acid addition salts are present in an amount of from about 25 mg. to about 100	00
10	EXAMPLE 9 dl - 2 - Amino - 1 - (3,4 - methylene -	9. d - 2 - Amino - 1 - (3.4 - methylene	90
e	Hydrochloride - 30 mg. Lactose 225 mg.	addition salts. 10. l - 2 - Amino - 1 - (3.4 - methylene -	95
)	Starch - 45 mg. The ingredients are mixed, granulated and compressed into a scored tablet which may be broken for divided doses if desired.	dioxyphenyl)-propane or its non-toxic acid addition salts, HASELTINE, LAKE & CO., 28, Southampton Buildings, London, W.C.2,	

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